WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: WO 92/18110 (11) International Publication Number: **A1** A61K 9/72 (43) International Publication Date: 29 October 1992 (29.10.92) (74) Agents: DANIELSSON, Sten et al.; AB Astra, Patent De-PCT/SE92/00186 (21) International Application Number: partment, S-151 85 Södertälje (SE). (22) International Filing Date: 24 March 1992 (24.03.92) (81) Designated States: AT, AT (European patent), AU, BB, BE (30) Priority data: 9101090-0 11 April 1991 (11.04.91) SE

(71) Applicant (for all designated States except US): AKTIEBO-LAGET ASTRA [SE/SE]; S-151 85 Södertälje (SE).

(72) Inventors; and (75) Inventors, and (75) Inventors, Applicants (for US only): TROFAST, Jan [SE/SE]; TROFAST, Eva [SE/SE]; Vapenkroken 34, S-224 47 Lund (SE). BYSTRÖM, Katarina [SE/SE]; Stora Vänern, Kullavägen, S-240 13 Genarp (SE). JAKUPOV-IC, Edib [YU/SE]; Smultronvägen 7, S-155 00 Nykvarn (European patent), BF (OAPI patent), BG, BJ (OAPI patent), BR, CA, CF (OAPI patent), CG (OAPI patent), CH, CH (European patent), CI (OAPI patent), CM (OAPI patent), CS, DE, DE (European patent), DK,

(OAPI patent), CS, DE, DE (European patent), DK, DK (European patent), ES, ES (European patent), FI, FR (European patent), GA (OAPI patent), GB, GB (European patent), GN (OAPI patent), GR (European patent), HU, IT (European patent), JP, KP, KR, LK, LU, LU (European patent), MC (European patent), MG, ML (OAPI patent), MN, MR (OAPI patent), MW, NL, NL (European patent), NO, PL, RO, RU, SD, SE, SE (European patent), SN (OAPI patent), TD (OAPI patent), TG pean patent), SN (OAPI patent), TD (OAPI patent), TG

Published

With international search report.

(OAPI patent), US.

(54) Title: PROCESS FOR CONDITIONING OF WATER-SOLUBLE SUBSTANCES

(57) Abstract

A process for providing water-soluble micronized substances, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such substances, which process is carried out by: a) reducing, if necessary, the residual water from the micronized substance by drying optionally at an elevated temperature and/or vacuum; b) conditioning said dried, micronized substances with a solvent; and c) eliminating residual solvent by storing in a dry place like vacuum or by purging with an inert gas.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT AU BB BE BF BG BJ CA CF CG CH CI CM CS DE DK	Austria Australia Barbados Belgium Burkina Faso Bulgaria Benin Brazil Canada Central African Republic Congo Switzerland Côte d'Ivoire Cameroon Czechoslovakia Germany Denmark	ES FI FR GA GB GN GR HU IT JP KP KR LI LK LU MC	Spain Finland France Gabon United Kingdom Guinea Greece Hungary Italy Japan Democratic People's Republic of Korea Republic of Korea Liechtenstein Sri Lanka I uxembourg Monaco	MG MI. MN MR MW NL NO PL RO RU SD SE SN SU TD TG US	Madagascar Mali Mongolia Mauritania Malawi Netherlands Norway Poland Romania Russian Federation Sudan Sweden Senegal Soviet Union Chad Togo United States of America
---	---	---	--	---	--

WO 92/18110 PCT/SE92/00186

PROCESS FOR CONDITIONING OF WATER-SOLUBLE SUBSTANCES

5 Field of the invention

The present invention relates to a process for providing water-soluble micronized substances, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such substances and which have improved physicochemical properties in the dry state, thereby facilitating the technical handling and significantly increasing the medical value of the substances.

15 Background of the invention

During the past few years, there have been frequent demonstrations of the fact that the appropriate selection of the most suitable crystalline modification significantly 20 can influence the clinical results of a given chemical entity. The chemical and physical stability of a solid compound in a particular dosage form can be modified by presenting the substance in the appropriate crystal form. Little information is available on the role of polymorphism 25 and crystal habit in solid dosage form and powder technology. It is, however, apparent that the appropriate selection of the most suitable crystalline modification, whether arising from polymorphic differences or as a result of solvate complex formation of both water-soluble substances 30 and less water-soluble substances, such as theophylline, often significantly can increase the medical value of a given drug in a particular dosage form. There are only a few statements available to predict the outcome of a crystallization procedure if e.g. the substance could be 35 involved in different polymorphic or pseudopolymorphic forms. Solid-state transformations may also occur during mechanical treatment, e.g. micronization and by pressure during tableting. While a few generalizations can be made concerning the influence of structural modifications on

WO 92/18110 PCT/SE92/00186

the tendency of a chosen compound to exhibit polymorphism or other phenomena, a complete understanding of this problem awaits further research. Often "trial and error" approaches are used to develop a successful formulation of a drug. It is necessary to establish the conditions whereby different forms of a substance might be converted to a single form thus eliminating differences in solid-state properties and subsequent different physico-chemical properties.

10

5

E. Shefter and T. Higuchi have measured the relative rates of dissolution of several crystalline solvated and non-solvated forms of important pharmaceuticals, J. Pharm.

15 Sci., 52 (8), (1963), 781-91.

L. van Campen, G. Zografi and J.T. Carstensen give in a review article an approach to the evaluation of hygroscopicity for pharmaceutical solids, Int. J. Pharmaceut. 5,

20 (1980), 1-18.

C. Ahlneck and G. Zografi describe the molecular basis of moisture on the physical and chemical stability of drugs in the solid state, Int. J. Pharmceut., 62, (1990), 87-95.

25

M. Otsuka et al. have calculated hydration data using various solid-state kinetic models for theophylline anhydrate powder, J. Pharm. Pharmacol., 42, (1990), 606-610.

30

Hak-Kim Chan and Igor Gonda have examined the properties of respirable crystals of cromoglycic acid by using different methods, J. Pharm. Sci., 78 (2), (1989), 176-80.

A more comprehensive discussion of factors relating to pharmaceutical preformulations and the physicochemical properties of drug substances is given by J.I. Wells in Pharmaceutical Preformulation: The Physicochemical

Properties of Drug Substances, John Wiley & Sons, New York (1988). See particularly the chapter about polymorphism pp 86-91.

5

Brief description of the invention

The object of the invention is to provide a process for water-soluble micronized substances, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such substances, whereby reducing the residual water from the micronized substances, conditioning said dried, micronized substances with a solvent and finally eliminating residual solvent from the substances.

Detailed description of the invention

- The object of the present invention is to provide a reliable process, where the desired polymorphic form can be conveniently and reproducibly prepared. The invention relates to a three step procedure:
- a. reducing, if necessary, the residual water from the micronized substance by drying optionally at an elevated temperature and/or vacuum.
 - conditioning said dried micronized substance with a solvent, and
- c. eliminating the residual solvent by storing the substance in a dry place, such as vacuum, or by purging with an inert gas.

The solvents used in the conditioning step b) are organic alcohols, ketones, esters, acetonitrile and the like, most preferably lower alcohols like methanol, ethanol, n-propanol, isopropanol; lower ketones like acetone, methylethylketone; ethylacetate, preferably in the vapour phase.

WO 92/18110 PCT/SE92/00186

According to one preferred embodiment the conditioning step b) is carried out in an inert gas containing solvent vapour.

5

15

20

25

30

The inert gas used in step c) and optionally in step b) is preferably nitrogen.

The preferred substances on which the invention is to be applied are carbohydrates, amino acids and drugs.

Carbohydrates, such as lactose, glucose, fructose, galactose, trehalose, sucrose, maltose, xylitol, mannitol, myoinositol and the like, and amino acids, such as alanine, betaine and the like, are often used as additives in pharmaceutical compositions e.g. as additives in certain inhalation formulations.

Terbutaline sulfate, salbutamol sulfate, fenoterol hydrobromide and bambuterol hydrochloride are highly selective β_2 -adrenergic agonist having bronchospasmolytic effect and are effective in the treatment of reversible obstructive lung ailments of various genesis, particularly asthmatic conditions. Disodium chromoglycate (DSCG) has been used as a prophylactic agent in the treatment of allergic bronchial asthma for many years.

The invention will be described by using lactose, terbutaline sulfate and salbutamol sulfate as examples. The phenomena of solvate formation and polymorphism are well recognized in the literature in the preformulation studies in the development phase for new drugs in the solid state. e.g. the US Pharmacopoeia recognizes >90 drug hydrates!

35

Many substances exist in different polymorphs (pseudopolymorphs) and several metastabile solvates with variable composition and physical properties like bulk density and

hygroscopicity. Several transformations between these polymorphs may occur at different velocity. These effects are operating when crystalline substances have been activated by various processes such as grinding, freeze 5 drying, micronization or recrystallization to produce regions of partial amorphous structure. The substances often will be obtained in an amorphous state or a metastable crystalline form when spray drying, freeze drying, rapid solvent quenching or when using controlled precipita-10 tion where both crystalline and amorphous forms can be prepared. The use of an amorphous form or a metastable crystalline form is often limited due to its thermodynamic instability. It is therefore a desire to convert the amorphous form or the metastable crystalline formto the 15 more stable crystalline state. The present invention deals with such physical and chemical changes, or more importantly, to anticipate them and the means by which these solid-state phenomena can be handled.

After recrystallization (or after spray drying/freezedrying) the substance has to be micronized to the final particle size required for e.g. inhalation. The particles should be less than 100 μm and preferably less than 10 μm. For crystalline substances, the micronization step seems to give an amorphous outer layer of the particle making the particle more sensitive to moisture.

30

35

It is an object of this invention to be able to reliably provide a crystalline form of certain water-soluble substances, which can be produced, stored and used, while maintaining the aerodynamic properties and specifications (particle size, particle form, hygroscopicity etc) required for inhalation of such substances. The particle size of the micronized substances is identical before and after the conditioning step as measured by different instruments like Malvern Master Sizer, culter counter or a microscope.

The conditioning of the substance probably rearrange the

outer layer of the crystals or the amorphous substance giving a more stable and less hygroscopic product.

In some instances it has been possible to use infrared spectroscopy in order to study the conversion of an amorphous form or a partly crystalline form into a stable crystalline form. Other methods available include BET gas adsorption, X-ray powder diffraction, microcalorimetry and differential scanning calorimetry (DSC). We have found that BET gas adsorption and microcalorimetry being the best methods for distinguishing the different forms of the tested compounds.

٨

5.9

Test results

8.4

5

10

15

20

The surface area measured by determining the quantity of a gas (nitrogen) that adsorbs as a single layer of molecules, a monomolecular layer on a sample is formed (Flowsorb II 2300, Micromeritics Co, USA). Surface area after the sample has been standing in high humidity for 24 hrs.

Micronized substance Non-conditioned substance
Conditioned substance
(m²/g) (m²/g) (m²/g)

Terbutaline sulfate:

11 - 12.5 < 3 7 - 9
Salbutamol sulfate:

With the low surface area, obtained when micronized substance has been stored at high humidity, the bulk substance has a great tendency to aggregate when stored, which make the substance very difficult for technical handling in

3

5

10

15

manufacturing the different formulations needed.

The interactions between certain substances and water vapour have also been studied by microcalorimetry. When said substances are subjected to water in the vapour phase they give off heat in a highly cooperative process. This moisture induced phase transition is however not observed for the conditioned substance. Thus, the conditioning process transforms the substance into a more stable form that is less sensitive to humidity.

Comparison of the heat given off by non-conditioned and conditioned substances when subjected to water vapour. Experiments are performed by a Thermal Activity Monitor 2277 (Thermometrics, Sweden).

Heat (J/g)

Relative humidity (%) Non-conditioned substance Conditioned substance

20

30

Terbutaline sulfate

58	3.6	0.1
7 5	6.2	0.1

25 Salbutamol sulfate

75 6 - 8 0.1

When spray-dried lactose has been conditioned in ethanol vapour for 100 hours at room temperature the energy given off was < 0.1 J/g, while the unconditioned lactose loses 40-44 J/g when subjected to water vapour.

The stability of the particles being conditioned were astonishing and will in a remarkable way increase the flexibility of the use of the substance for different formulations.

Experimental procedure

The invention is further illustrated but not limited by the following example.

5

10

25

Example 1

3.6 kg terbutaline sulphate micronized was dried in a stainless steel column with 200 mm diameter at 90°C in vacuum for 23 hours. The dried substance was cooled to about 30°C and the pressure was normalized with ethanolsaturated nitrogen gas. 70 ml/min of ethanol-saturated nitrogen gas was then passed through the 200 mm diameter column for 60 hours to condition the substance. During this time the column was inverted a few times. The residual 15 solvent was eliminated by purging with nitrogen gas for 2 hours and the product, about 3.5 kg, was packed in double plastic bags with a drying agent between the bags.

20 Example 2

In one experiment 1 g micronized salbutamol sulfate was kept at room temperature for 24 hours in a closed vessel containing a beaker filled with ethanol. The sample was removed and stored in a completely dry environment over night in order to eliminate traces of ethanol. The sample was subjected for analysis (see test results given above).

It is necessary to introduce stirring or tumbling of the substance when conditioning in larger scale. 30

Example 3

1 g spray-dried amorphous lactose was treated as in exemple 2. The time kept in the saturated ethanol vapour was 100 35 hours. After removal of residual ethanol, the sample was subjected for calorimetric analysis (see test results given above).

CLAIMS

- 5 1. A process for providing water-soluble micronized substances, which can be produced, stored and used while maintaining the aerodynamic properties required for inhalation of such substances, c h a r a c t e r i z e d in
- a) reducing, if necessary, the residual water from the
 micronized substance by drying optionally at an elevated
 temperature and/or vacuum,
 - b) conditioning said dried, micronized substances with a solvent, and
- c) eliminating residual solvent by storing in a dry place is like vacuum or by purging with an inert gas.
- A process according to claim 1, c h a r a c t e r i z e d in that the solvent used in the conditioning step b) is ethanol, acetone or the like, preferably in the vapour phase.
 - 3. A process according to claim 2, c h a r a c t e r i z e d in that the solvent used in step b) is ethanol.
- 25 4. A process according to any one of claims 1-3, c h a r a c t e r i z e d in that the conditioning step b) is carried out in an inert gas containing solvent vapour.
- 5. A process according to any one of claims 1-4,
 30 characterized in that the inert gas used in step c) and optionally in step b) is nitrogen.
- 6. A process according to any one of claims 1-5, characterized in that the substances are additives, such as carbohydrates and amino acids.
 - 7. A process according to claim 6, c h a r a c t e r i z e d in that the carbohydrates used are lactose,

glucose, fructose, galactose, trehalose, sucrose, maltose, xylitol, mannitol, myoinositol or the like and the amino acids used are alanine, betaine or the like.

- 5 8. A process according to any one of claims 1-6, characterized in that the substances are drugs.
- A process according to claim 8, c h a r a c t e r i z e d in that said drugs are antiasthmatic or antiallergic substances.
- 10. A process according to claim 8, c h a r a c t e r i z e d in that said antiasthmatic or antiallergic
 15 substances are selected from terbutaline sulfate, salbutamol sulfate, fenoterol hydrobromide, bambuterol hydrochloride, terfenadine and disodium chromoglycate.

INTERNATIONAL SEARCH REPORT

International Application No PCT/SE 92/00186

I. CLASSIFICATIO	N OF SUBJECT MATTER (if several classifi	ication symbols apply, indicate all) ⁶	
According to Interna IPC5: A 61 K	ational Patent Classification (IPC) or to both N 9/72	ational Classification and IPC	
II. FIELDS SEARCH		7	
Oleralisation System	Minimum Docume	ntation Searched Classification Symbols	
Classification System		riassineation cymbols	
IPC5	A 61 K		*
		than Minimum Documentation s are included in Fields Searched ⁸	
SE,DK,FI,NO	classes as above		
III. DOCUMENTS C	ONSIDERED TO BE RELEVANT ⁹		
Category * Citat	ion of Document, ¹¹ with indication, where app	propriate, of the relevant passages ¹²	Relevant to Claim No. ¹³
10	l, 0436110 (BIOCHEMIE GESE D July 1991, age 3, line 17 - line 54	LLSCHAFT M.B.H.)	1-10
	, 4405598 (BROWN K ET AL.) ol. 4, line 4 - line 27 	20 September 1983,	1-10
33	1, 8607547 (GERGELY G. ET AL) 1 December 1986, ee the whole document		1-10
"A" document defi considered to	ies of cited documents: ¹⁰ ning the general state of the art which is not be of particular relevance	"T" later document published after or priority date and not in conflicited to understand the principle invention	the international filing date ict with the application but e or theory underlying the
"E" earlier docum filing date	ent but published on or after the international	"X" document of particular relevance cannot be considered novel or o	e, the claimed invention
"L" document whi which is cited	ch may throw doubts on priority claim(s) or to establish the publication date of another	involve an inventive step "Y" document of particular relevance	
citation or oth "O" document refe other means	er special reason (as specified) erring to an oral disclosure, use, exhibition or	cannot be considered to involve document is combined with one ments, such combination being in the art.	or more other such docu-
"P" document pub tater than the	lished prior to the international filing date bu priority date claimed	t "&" document member of the same	patent family
IV. CERTIFICATION		Date of Mailing of this International S	earch Ronort
16th July 199	mpletion of the International Search 92	1992 -07- 2 1	uar ori inapori
International Searchi	ng Authority	Signature of Authorized Officer association of Millianl G	Kegt:
	DISH PATENT OFFICE	Anneli Jönsson	0

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.PCT/SE 92/00186

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report.

The members are as contained in the Swedish Patent Office EDP file on

The Swedish Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

EP-A1- 0436110 91-07-10 US-A- 4405598 83-09-20	AU-D- CA-A- JP-A- WO-A- CA-A- AU-B- AU-D- BE-A- CA-A- DE-A-C- FR-A-B- GB-A- JP-C- JP-A- JP-B- LU-A-	6870191 2030581 3172181 91/07948 	91-06-26 91-05-25 91-07-25 91-06-13 82-03-23 80-10-16 78-08-03 77-07-25 84-10-16 77-08-04 77-08-26 80-03-19 87-02-26 77-08-09
US-A- 4405598 83-09-20	AU-B- AU-D- BE-A- CA-A- DE-A-C- FR-A-B- GB-A- JP-C- JP-A- JP-B-	512593 2162677 850727 1176171 2703119 2339604 1562901 1366290 52094411	80-10-16 78-08-03 77-07-25 84-10-16 77-08-04 77-08-26 80-03-19 87-02-26 77-08-09
	NL-A- SE-B-C- SE-B-C- SE-A- SE-A- AU-D- BE-A- CH-A- DE-A-C- FR-A-B- GB-A-B- JP-B- JP-C- JP-A- NL-A- SE-B-C- SE-A-	76661 7700911 442267 442268 7700888 8107278 522792 3805778 869055 1112567 627075 2831419 2397833 2001334 1011615 1532970 54035209 7807625 443087 7807934	86-07-10 77-08-03 77-08-02 85-12-16 85-12-16 77-07-31 81-12-04 82-06-24 80-01-17 79-01-17 81-11-17 81-12-31 79-02-01 79-02-16 79-01-31 89-02-27 89-11-24 79-03-15 79-01-23 86-02-17 79-01-20
WO-A1- 8607547 86-12-31	EP-A-B- JP-T- US-A- US-A-	0258258 63501137 4876802 4911930	88-03-09 88-04-28 89-10-31 90-03-27